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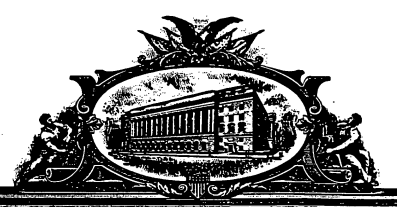
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By Authority of the COMMISSIONER OF PATENTS AND TRADEMARKS

H. PHILLIPS
Certifying Officer

Wowk

[45] Sept. 11, 1973

[54]	BIS[HYDROCARBYL(HALO)(MERCAPTO)- TIN]OXIDE				
[75]	inventor:	Anatole Wowk, Edison, N.J.			
[73]	Assignee:	M & T Chemicals Inc., New York, N.Y.			
[22]	Filed:	June 24, 1971			
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	Rela	ted U.S. Application Data			
[62]	Division of	Ser. No. 43,231, June 3, 1970, abandoned.			
[52]	U.S. CL	260/429.7, 260/45.75 K			
[51]	Int. Cl				
[58]		earch 260/429.7			
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Primary Examiner—Werten F. W. Bellamy Attorney—Lewis C. Brown et al.

57] ABSTRACT

This invention comprises novel compounds of the f r-mula:

wherein R and R' are hydrocarbons and X is halogen, methods of preparing these novel compounds, and polymers stabilized by these novel compounds against the deteriorative effects of heat and light.

8 Claims, No Drawings

BIS[HYDROCARBYL(HALO)(MERCAPTO)TIN-]OXIDE

This application is a divisional application of Ser. N 43,231, filed June 3, 1970, and now abandoned.

This invention relates to novel organotin compounds and to novel stabilized halogen-containing polymer compositions.

Halogen-containing polymers, including homopolymers and copolymers of vinyl chloride and vinylidene 1 chloride, are materials which have proven useful, because of their desirable physical properties. Uses of these materials have been limited, however, by their inherent instability when exposed to conditions of heat and light. Under such conditions, the halogen- 15 containing polymer may discolor, become brittle, crack, check or otherwise suffer deterioration of its physical properties. Thus, unstabilized chlorine-containing polymer compositions may be highly unsatisfactory for outdoor use where they may be subjected to both heat 20 and light. Various techniques are known for stabilizing these polymers against degradation in the presence of heat alone or light alone but no single stabilizer compound has been found which confers the high degree of both heat and light stability which is required for out- 25

A further defect of prior art stabilizers is that they may normally be viscous liquids or pasty gels. They may thus be considerably more difficult to handle than solid stabilizers.

It is an object of this invention to provide novel organotin compounds which may be characterized by their unexpectedly superior physical properties. It is a further object of this invention to provide a novel chlorine-containing polymer compositions characterized by their high resistance to degradation during outdoor exposure.

This invention also comprises halogen-containing polymers stabilized against the deteriorative effects of heat and light comprising a halogen-containing polymer and a stabilizing amount of a compound of the formula:

wherein R and R' are hydrocarbons and X is halogen. The novel compounds of this invention may be prepared be reacting a compound of the formula RSn(OH), X wherein R is a hydrocarbon radical selected from the group consisting of alkyl, alkenyl, cycloalkyl, aralkyl, aryl, and alkaryl and X is halide with one or more primary mercaptans, secondary mercaptans, tertiary mercaptans, mercapto alcoh is, esters of alcohols, m reapto acids or, mercapto acid esters.

The novel compounds f this invention thus include those of the formula:

In these f rmulae R and R' are hydrocarbon radicals selected from the group consisting falkyl, cycloalkyl, aralkyl, alkaryl, and aryl. R' may be selected from the same group as R. R' may also be a residue of a carboxylic acid ester —R"COOR".

In these formulae R" represents a methylene or an ethylene radical and R" is selected from the same group as R. R and R' are hydrocarbon radicals selected from the group consisting of alkyl, cycloalkyl, aralkyl, alkaryl, and aryl, and X is a halogen atom selected from the group consisting of chlorine, bromine and iodin . When R or R' is alkyl, it may be a straight chain alkyl or a branched alkyl, including methyl, ethyl, n-propyl, isopropyl, n-butyl, isobutyl, sec-butyl, tert-butyl, namyl, neopentyl, isoamyl, n-hexyl, isohexyl, heptyls, octyls, decyls, dodecyls, tetradecyl, octadecyl, etc. Preferred alkyl includes alkyls having less than about 20 carbon atoms. When R or R' is cycloalkyl, it may typically be cyclopentyl, cyclohexyl, cycloheptyl, cyclooctyl, etc. When R or R' is aralkyl, it may typically be benzyl, β -phenylethyl, γ -phenylpropyl, β -phenylpropyl, etc. When R or R' is aryl, it may typically be phenyl, naphthyl, etc. When R or R' is alkaryl, it may typically be tolyl, xylyl, p-ethylphenyl, p-nonylphenyl, etc. R or R' may be inertly substituted, e.g. may bear a nonreactive substituent such as alkyl, aryl, cycloalkyl, aralkyl, alkaryl, ether, etc.

Typical novel compounds of this invention include bis(monobutylmonochlorotin lauryl mercaptide) oxide, bis(monobutylmonochlorotin octyl mercaptide) oxide, bis(monobutylmonochlorotin benzyl mercaptide) oxide, bis(monobutylmonochlorotin xylyi mercaptide) oxide, bis(monobutylmonochlorotin cyclohexyl mercaptide) oxide, bis(monobutylmonochlorotin phenyl mercaptide) oxide, bis(monooctylmonochlorotin lauryl mercaptide) oxide, bis(monooctylmonochlorotin octyl mercaptide) oxide, bis(monooctylmonochlorotin benzyl mercaptide) oxide, bis(monooctylmonochlorotin oxid . mercaptide) xylyl bis(monooctylmonochlorotin cyclohexyl mercaptide) oxide, bis(monocyclohexylmonochlorotin lauryl mercaptide) oxide, bis(monomethylmonochlorotin lauryl mercaptide) oxide, bis(monomethylmonochlorotin benzyl mercaptide) oxide, bis(monomethylmonochlorotin cyclohexyl mercaptide) oxide, bis(monomethylmonochlorotin xylyl dimercaptide) oxide, bis(monobutylmonochlorotin isooctyl mercaptoacetate) oxide, bis(monooctylmonochlorotin isooctyl mercaptoacetate) oxide, bis(monocyclohexylmonochlorotin isooctyl mercaptoacetate) oxide, bis(monomethylmonochlorotin isooctyl mercaptoacetate) oxide, bis(monophenylmonochlorotin isooctyl mercaptoacetate) oxide, bis(monobutylmonochlorotin lauryl mercaptoacetate) oxide, bis(monobutylmonochlorotin benzyl mercaptoacetate) oxide, bis (monobutylmonoisooctyl mercaptopropionate) chlorotin bis(monooctylmonochlorotin isooctyl mercaptopropionate) oxide, bis(monocycloh xylm nochlorotin isooctyl mercaptopropi nate) xide, bis(mon methylm nochlorotin isooctyl mercaptopropionate) oxide, bis)monophenylmenochlorotin isooctyl mercaptopropionate) oxide, bis(mon butylmonochlorotin lauryl mercaptopropionate) oxide, bis(monobutylmonochlorotin benzyl mercaptopropionate) oxide, bis(m nobutylmonochlorotin isooctyl mercaptopropionate) oxide, and bis(monobutylmonochlorotin dodecyl mercaptopropionate) oxide.

The mono-halo-mono organotin oxide reactant [RSnX], I this invention is prepared from the compound RSnX, Typical c mpounds RSnX, include the following: ethyltin trichloride, propyltin trichloride, n-butyltin trichloride, hexyltin trichloride, octyltin trichloride, benzyltin trichloride, benzyltin trichloride, butyltin trichloride, ethynyltin trichloride, butyltin tribromide, and n-octyltin tribromide.

Compounds such as butyltin bromide dichloride may 10 be employed. Preferably the compound RSaXa is a chloride; and the preferred compounds may be nbutyltin trichloride and n-octyltin trichloride. The process for preparing the compound RSn(OH), X includes maintaining an aqueous reaction mixture containing RSnX₂; adding to said reaction mixture a catalytic amount of a catalyst selected from the group consisting of aluminum chloride, titanium tetrachloride, bismuth trichloride, ferric chloride, cobalt chloride, nickel chloride, cadmium chloride, zirconium tetrachloride, boron trifluoride etherate, mercuric chloride, cupric chloride, trifluoracetic acid, and zinc fluoride; thereby forming a precipitate containing RSn(OH),X; and recovering said precipitate from said aqueous reaction mixture. In practice, 10 - 200 parts, say 17.5 parts of RSnX, are added to 100 parts of aqueous reaction medium, preferably water. Catalyst, preferably aluminum chloride. is added in an amount of 0.1 - 10 parts, say 0.95 parts, and the solution allowed to stand for up to 2-3 hours. 30 The solid precipitate which forms may be separated as by filtration and washed with 10 - 1,000 parts, say 15 parts of water. After further washing with 6 - 60 parts, say 12 parts of e.g. acetone, the precipitate may be airdried. The product, typically obtained in an amount of 35 2.5 parts, is recovered from the filtrate.

The process for preparing RSn(OH)₂X when R is alkyl includes maintaining an aqueous reaction mixture containing RSnX_a; heating said reaction mixture thereby forming a precipitate containing RSn(OH)₂X 40 product; and recovering said precipitate from said aqueous reaction medium.

Typical reactants [RSnXO], include the following: ethyltin chloride oxide polymer, propyltin chloride oxide polymer, n-butyltin chloride oxide polymer, iso-45 butyltin chloride oxide polymer, hexyltin chloride oxide polymer, phenyltin chloride oxide polymer, o-tolyltin chloride oxide polymer, benzyltin chloride oxide polymer, butenyltin chloride oxide polymer, butenyltin chloride oxide polymer, and ethynyltin chloride oxide polymer; ethyltin chloride dihydroxide, propyltin chloride dihydroxide, iso-butyltin chloride dihydroxide, hexyltin chloride dihydroxide, octyltin chloride dihydroxide, phenyltin chloride dihydroxide, benzyltin chloride dihydroxide, benzyltin chloride dihydroxide, butenyltin chloride dihydroxide, butenyltin chloride dihydroxide.

A reaction mechanism for the f rmati n of the novel compounds f this inventi n is clearly illustrated by the following equation:

99 **- 2000**

 $[B8nX0]_a + nH8R \longrightarrow \frac{n}{2} \begin{bmatrix} B8 \\ X \end{bmatrix}_0 + nH_a$

These equations graphically show that a reaction ccurs between the m reapto radical and the compound RSn(OH)₂X whereby water splits ff and a chemical bond is formed between the tin and sulfur. These equations further illustrate the necessity of two molecules f the mercapto compound per mole of the compound RSn(OH)₂X to obtain the desired reaction and the desired novel compound having a 2:1 ratio of sulfur to tin.

It is within the scope of this invention that pure, im10 pure or commercial grades of the reactants may be employed satisfactorily. In general, pure compounds of
the above formulae may be prepared from pure raw
materials. However, these novel compounds may be diluted with innocuous, inert materials thereby permitting the use of technical grades of materials or intermediates in their preparation.

Any suitable reaction temperatures may be employed. It is ordinarily preferred to use room or slightly elevated temperatures of the order up to about 100°C. The exothermic nature of the reaction between the compound RSn(OH)₂X and the sulfur containing compounds affords a considerable saving in the expenditure of external heat. The presence of water produced by the condensation reaction usually requires an additional heating or refluxing in order to strip said water from the reaction product, though it may be removed in any suitable matter.

The use of inert organic solvents as a medium for a reaction such as toluene, benzene, methyl alcohol, etc., is contemplated. The presence of such solvent facilitates the desired reaction. The solvent may be eliminated from the reaction product at the completion of the reaction by any suitable means. This may be accomplished by vaporizing the solvent under vacuum at elevated temperatures. Pressures of about 2 to 30 mm of mercury at temperatures of 75°C to 160°C are satisfactory in affecting the removal of toluene or like solvent from the reaction product.

By these procedures, the novel compounds of this invention can be obtained in almost theoretical yields. These yields are indicative that it is not necessary to use an excess of either reactant, the total amount of starting materials being substantially utilized in the formation of the final reaction product.

Polymers or resins which may be stabilized by practice of this invention are halogen-containing organic polymers typically those which contain chlorine atoms bonded to the polymer chain. These polymers include polyvinyl chloride-type polymers, e.g. polyvinyl chloride, polyvinylidene chloride, etc. They may also include copolymers formed by the copolymerization of vinyl chloride or vinylidene chloride with each other or with other ethylenically unsaturated monomers. Ethylenically unsaturated monomers may be compounds which contain polymerizable carbon-to-carbon double bonds and may include acrylates such as acrylic acid, ethyl acrylate, acrylonitrile, etc.; vinyl monomers such as styrene, vinyl acetate, etc.; maleates such as maleic acid, maleic anhydrid, maleate esters, tc.

acid, maleic anhydrid, maleate esters, tc.

The polymers may be either "rigid" r "flexible".

When "rigid" polym rs are employed, they may include impact modifiers, pigm nts and/r fillers, lubricants, etc., in addition to the resin and stabilizer.

When "flexible" polymers are employed, they may include plasticizer (primary and secondary), pigments and/or fillers, lubricants, etc., in addition to the resin and stabilizer.

In general the synthetic resins which can be stabilized according to this invention include the following polymers which may or may not be mixed with other stabilizers, additives, flameproofing agents, dyes, pigments, etć.:

a. Homopolymer of vinyl chloride.

b. Homopolymer of vinylidene chloride,

- c. Copolymens of vinyl chloride and acrylonitrile,
- d. Copolymers of vinylidene chloride and acryloni-
- e. Copolymers of vinylidene chloride, acrylonitrile and N-isopropylacrylamide,
 - f. Copolymers of vinyl chloride and vinyl acetate,
- g. Copolymers of vinyl chloride, acrylonitrile, and Nbutvrvlacrylamide.
- h. Copolymers of vinyl chloride, methyl methacrylate and vinyl acetate,
- i. Copolymers of vinyl chloride or vinylidene chloride with acrylonitrile and N-vinylpyridine,
- with acrylonitrile mixed with a homopolymer of an Nalkylacrylamide (e.g. N-isopropylacrylamide, Noctylacrylamide, etc.),
- k. Copolymers of vinyl chloride or vinylidene chloride with acrylontrile mixed with a homopolymer or co- 2^{4} polymer of a-vinylpyrrolidone,
- 1. Copolymers of vinyl chloride or vinylidene copolymers of an N-alkylacrylamide and acrylonitrile.
- m. Other copolymers and mixtures of homopolymers or copolymers of vinyl chloride and/or vinylidene chloride with or without other monoolefinically unsaturated polymerizable monomers containing from 2 to 20 or more carbon atoms, especially those containing from 2 to 8 carbon atoms and no more than one nitrogen atom and no more than 2 oxygen atoms.

The methods for preparing the synthetic resins described above are well known and such methods and a great variety of such synthetic resins are described in the prior art. These synthetic resins are useful in preparing fibers, films, molding compositions, coating materials, wrapping materials, electrical insulation, fabrics, rope, plastic, pipe, paints, laminating materials for safety glass, adhesives, etc. Synthetic resins as stabilized in accordance with this invention are especially advantageous in synthetic fibers and products made therefrom such as rugs, wearing apparel, draperies, seat covers, upholstery, rope, cigarette filters, etc. Resistance to weathering (e.g. moisture and ultraviolet light) is especially important for items exposed to out of doors conditions. Resistance to ultraviolet light is also important for rugs, draperies, automobile seat covers, porch furniture upholstery and the like which may receive considerable sunlight.

The following examples are for illustration and are not to be construed as limiting the scope of the invention.

EXAMPLE 1

Preparation f bis(monobutylmonochl rotin isooctyl mercaptopropionate) oxid

Bis(monobutylmonochlorotin isooctyl mercaptopropionate) oxide was prepared by adding 22.7 grams (0.10 moles) of butylchlorotin oxide polymer and 21.8 grams (0.10 moles) of isooctyl mercaptopropionate to 150 milliliters f benzene contained in a reaction vessel equipped with a stirrer, a th rmometer and a reflux condenser ab v a Dean-stark water trap. The water

was azeotroped out at 80°C to 82.5°C. The benzene solvent was stripped leaving the product. The product exhibited a weight of 44.5 grams. Analysis f the product indicated the following:

Elemental Percentage		
of .	Calculated	Found
Se .	27.20	26.99
8	7.34	7.30

EXAMPLE 2

Preparation of bis(monobutylmonochloro-Shydroxyethylmercaptotin) oxide

Bis(monobutylmonochloro-S-hydroxyethylmercaptotin) oxide was prepared by adding 18.4 grams (0.075 moles) of butylchlorotin dihydroxide and 5.9 grams (0.075 moles) of 2-mercaptoethanol to 120 milliliters of benzene in a 200 milliliter reaction flask equipped with a stirrer, a thermometer, and a reflux condenser. j. Copolymers of vinyl chloride or vinylidene chloride 20 The foregoing procedure was followed. The resulting yellow liquid product exhibited a weight of 21.7 grams and was analyzed as containing the following:

	Elemental Percentage	•		
	of C.	Calculated	•	Found
3	S S	40.02 10.81		40.97 10.03
	ä	11.95		14,34

This invention is also of a novel stabilizing composition for stabilizing a vinyl chloride homopolymer or copolymer against the deteriorative effects of heat which comprises as a first stabilizer a novel compound of this invention exhibiting a compound of the formula

wherein R and R' are hydrocarbons and X is halogen and as a second stabilizer an organotin compound exhibiting two direct carbon to tin bonds and two direct sulfur to tin bonds.

Among the second stabilizers operable in the practice of this invention are compounds of the formula R's Sn(SR")₂. In the formula R and R' are hydrocarbon radicals selected from the group consisting of alkyl, cycloalkyl, aralkyl, alkaryl, and aryl. R" may be selected from the same group as R and R'. R" may also be a residue of a carboxylic acid ester -R""COOR"". In these formulae R"", R""" (which may be the same as each other or different) may be selected from the same group as R and R' are selected from. Typical second stabilizers include dioctyltin bis(isooctyl mercaptoacctate) and dibutyltin bis(lauryl:mercaptide).

Typical preferred second stabilizers include:

dibutyltin bis(lauryl mercaptide)

dibutyltin bis(octyl mercaptide) dibutyltin bis(benzyl mercaptide)

dibutyltin bis(xylyl m reaptide)

dibutyltin bis(cyclohexyl mercaptide)

dibutyltin bis(phenyl mercaptide)

dioctyltin bis(lauryl mercaptide)

dioctyltin bis(octyl mercaptide) dioctyltin bis(benzyl mercaptid)

dioctyltin bis(xylyl mercaptide) dioctyltin bis(cycl hexyl mercaptide) dicyclohexyltin bis(lauryl mercaptide)
dimethyltin bis(lauryl mercaptide)
dim thyltin bis(benzyl mercaptide)
dimethyltin bis(cyclohexyl mercaptide)
dimethyltin bis(sylyl dimercaptide)
dibutyltin bis(isooctyl mercaptoacetate)
dioctyltin bis(isooctyl mercaptoacetate)
dicyclohexyltin bis(isooctyl mercaptoacetate)
dimethyltin bis(isooctyl mercaptoacetate)
diphenyltin bis(isooctyl mercaptoacetate)
dibutyltin bis(lauryl mercaptoacetate)
dibutyltin bis(benzyl mercaptoacetate)
dibutyltin bis(benzyl mercaptoacetate)

Other preferred second stabilizers include dibutyltin bis(cyclohexyl maleate), dioctyltin bis(cyclohexyl maleate), dibutyltin bis(2-methylcyclohexyl maleate), diamyltin bis(4-ethylcyclohexyl maleate), dibutyltin bis(4-methylcyclohexyl maleate), dibutyltin bis(2-cyclohexylcyclohexyl maleate), dibutyltin bis(4-cyclohexylcyclohexyl maleate), dibutyltin bis(4-cyclohexylcyclohexyl maleate), dibutyltin bis(2-ethynylcyclohexylmaleate), dibutyltin bis(4-hydroxymethylcyclohexylmaleate), dibutyltin bis(2,6-dimethylcyclohexylmaleate), dibutyltin bis(4,6-trimethylcyclohexylmaleate), dibutyltin bis(4-secbutylcyclohexylmaleate), dioctyltin bis(4-tert-butylcyclohexylmaleate), etc.

The preferred second stabilizer is dibutyltin bis (isooctyl mercaptoacetate).

The first and second stabilizers may be present in stabilizing amounts. Typically the first stabilizer is used in an amount of 0.05 to 10 parts by weight of 100 parts by weight of halogen-containing, typically vinyl chloride resins. Preferably, it may be used in the amount of 0.25 - 2 parts, preferably 0.3 parts per 100 parts of halogen-containing resin.

Typically the second stabilizer is present in an ³⁵ amount of 0.1 – 10 parts per 100 parts by weight of halogen-containing resin. Preferably the amount used may be 1, say 2 parts per 100 parts of halogen-containing resin. Preferably the second stabilizer is present in an amount about five times the amount of the first stabilizer. Preferred results achieved when the second stabilizer is present in an amount of at least about twice that of the first stabilizer.

Thus a preferred heat-stable vinyl chloride polymer composition of this invention comprises 100 parts by weight of a vinyl chloride polymer; a stabilizing amount, typically 0.05 – 10 parts of a first stabilizer having at least one direct halogen to tin bond and at least one carbon to tin bond wherein R is selected from the group consisting of alkyl having 1–20 carbon atoms, aryl, alkaryl, aralkyl, and cyclo-alkyl; and a second stabilizer R'₂Sn(SR'')₂ wherein R' is a hydrocarbon radical selected from the group consisting of alkyl, cycloalkyl, aralkyl, alkaryl, and aryl, R'' may be a residue of a carboxylic acid ester or a radical selected from the same group as R', and n is an integer 1–3.

The novel heat and light stable compositions of this invention may be f rmulated by such techniques as milling, dry blending, Banbury blending, or any oth r commonly empl yed formulating techniques.

On th formulating m thods which is particularly convenient is the formation f a stabilizer composition containing the first stabilizer and the second stabilizer. This stabilizer composition is later added to, and th roughly mixed with, the vinyl chl ride polymer. Where this technique is employed the stabilizer composition typically c mprises stabilizing am unts of the stabiliz-

rs, typically 0.05 to 5 parts by weight f the first stabilizer and 0.1 to 10 parts by weight of the second stabilizer. Preferably, it comprises about 0.05 to 2, say 0.3 parts of the first stabilizer and about 0.5-4, say 0.2 parts of the second stabilizer. Preferably the second stabilizer may be present in an amount greater than the amount of the first stabilizer.

Whatever formulating technique be employed, it will be desirable to substantially complete and uniformly 10 disperse the first stabilizer and the second stabilizer throughout the vinyl chloride polymer conposition.

It is a feature of the novel stabilizer systems of this invention that it permits attainment of stabilized halogen-containing polymers and resins, particularly vinyl halide polymers such as vinyl chloride characterized by their resistance to the deteriorative effect of heat. The degrees of stabilization attained in such systems may be considerably in excess of that previously attainable by any prior art stabilizer system.

Because of the outstanding properties of this novel stabilizer system, it is possible to effect stabilization with lower quantities and thereby to obtain a more effective system on a cost-performance basis.

In order to set forth clearly the novel features of this invention and to illustrate the unexpected and outstanding results which may be attained by practice of this invention, the following illustrative examples may be set forth wherein all parts are parts by weight unless otherwise indicated.

In these examples, the rigid vinyl chloride polymer employed was that having a specific gravity of 1.40, a Shore Durometer "D" hardness of 80 and an ultimate tensile strength of about 7,000 psi sold under the trademark "Geon 103 EP".

EXAMPLE 3

Testing of bis(monobutylmonochloro-S-hydroxyethylmercaptotin) oxide

The preferred illustrative first stabilizer employed was bis(monobutylmonochloro-S-hydroxyethylmer-captotin) oxide. The second stabilizer used was dibutyltin bis(isooctyl mercaptoacetate). 100 parts of the vinyl chloride polymer was tested with addition of the stabilizer combination of the following amounts:

vinyl chloride polymer first stabilizer dibutyltin bis(isooctyl snercaptoscetate) 100 part 0.2 "

. . .

The results of the heat stability tests are as follows: The foregoing composition was thoroughly blended by placing the polyvinyl chloride on a two-roller differential mill which was oil-heated to a temperature of 175°C together with the noted quantity of first and second stabilizers, and the mixture was milled for about 5 minutes. A continuous band of the composition formed around one f the rollers. This band was cut and the composition was removed from the hot roller as a continuous sheet. Squares f this material measuring 2:54 cm × 2.54 cm were cut f r heat stability testing.

For th heat stability test, the squares were placed in an air ov n regulated to maintain a temperature of 190°C. Samples f each composition were removed from the oven at 15 minute intervals and were rated visually as to color chang and degradation according to the f llowing scale:

7 — clear, water-white

6 - ff-white

5 - slightest degree f yellowing

4 — definite yellow color

3 - deep yellow-brown color

2 — deep brown color

1 - dark brown to black color

The length of time in minutes required to reach a value of 3 or less was recorded as the Heat Stability Value.

EXAMPLE 4 (control)

In control Example 4 the dibutyltin bis(isocctyl mercaptoacetate) was added in amount of 2 parts and no other stabilizer was present.

The results of the heat stability tests are as follows: 15

Example	Heat Stability Value in Minutes	Color After Milling	
3	75	7	
4	75 ·	7	

It will be apparent from an inspection of Examples 3 and 4 that the Heat Stability Value (H.S.V.) of the novel product of this invention is outstanding. The H.S.V. of this novel composition may be at least as long as 75 minutes. Furthermore, the Color After Milling of the novel product is 7, which (when considered together with the 75 minute H.S.V.) is outstandingly superior. Accordingly, it will be apparent to those skilled in the art that practice of the novel process of this invention permits attainment of polymeric resins characterized by extremely high degree of heat stability.

Although this invention has been illustrated by reference to specific examples, changes therein which clearly fall within the scope of the invention will be apparent to those skilled in the art. It is, therefore, to be 35 limited solely by the scope of the appended claims.

I claim:

1. A compound of the formula



wherein R is selected from the group consisting of alkyl radical containing between one and twenty carbon atoms, cycloalkyl, aryl, aralkyl and alkaryl radicals; R' represents a radical selected from the same group as R, a β-hydroxyethyl radical, hydroxypropyl radical or a radical of the formula —R"COOR" wherein R" represents a methylene or an ethylene radical and R" is selected from the same group as R and X is a halogen radical.

2. The compound claimed in claim 1 wherein R is alkyl.

3. The compound claimed in claim 1 wherein R is butyl.

4. The compound claimed in claim 1 wherein R is octvl.

5. The compound claimed in claim 1 wherein X is chloride.

6. The compound of claim 1 wherein R and R' are alkyl of from 1 to 20 carbon atoms.

7. The compound claimed in claim 1 wherein said compound is bis(monobutylmonochloro-Shydroxypropylmercaptotin) oxide.

8. The compound claimed in claim 1 wherein said compound is bis(monobutylmonochloro-S-hydroxyethylmercaptotin) oxide.

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4Ω

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